

Clinical Trial Protocol

Iranian Registry of Clinical Trials

21 Jul 2026

Comparison of the effect of intra tracheal injection of budesonide with surfactant and surfactant alone in prevention of Bronchopulmonary Dysplasia in preterm infant under ۳۰ week with Respiratory Distress Syndrome A Randomized Cilinical Trial

Protocol summary

Study aim

Determination of the effect of budesonide intratracheal injection in the prevention of bronchopulmonary dysplasia in premature infants with a birth age of less than 30 weeks of gestation and respiratory distress syndrome

Design

A clinical trial with a control group

Settings and conduct

Kamali Hospital

Participants/Inclusion and exclusion criteria

Premature infants less than 30 weeks of age with respiratory distress syndrome

Intervention groups

Preterm infants who meet the conditions for entering the study are included in the study as available sampling and based on the list Randomization falls into one of the following two groups: mg/kg intervention in the first group including: intratracheal injection of surfactant 200 and followed by mg/kg and surfactant 200 budesonide 0.25 mg/kg in the second intervention group including intratracheal injection Budesonide 0.25 mg/kg (0.5 ml/kg) along with 100 mg/kg (1.25 ml/kg if needed)

Main outcome variables

Examination in terms of oxygen dependence

General information

Reason for update

Acronym

IRCT registration information

IRCT registration number: **IRCT20221016056205N1**

Registration date: **2023-05-15, 1402/02/25**

Registration timing: **registered_while_recruiting**

Last update: **2023-05-15, 1402/02/25**

Update count: **0**

Registration date

2023-05-15, 1402/02/25

Registrant information

Name

Sima Kianpisheh

Name of organization / entity

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Recruitment status

Recruitment complete

Funding source

Expected recruitment start date

2022-11-22, 1401/09/01

Expected recruitment end date

2023-05-20, 1402/02/30

Actual recruitment start date

empty

Actual recruitment end date

empty

Trial completion date

empty

Scientific title

Comparison of the effect of intra tracheal injection of budesonide with surfactant and surfactant alone in prevention of Bronchopulmonary Dysplasia in preterm infant under ۳۰ week with Respiratory Distress Syndrome A Randomized Cilinical Trial

Public title

Comparison of the effect of intra tracheal injection of budesonide with surfactant and surfactant alone in prevention of Bronchopulmonary Dysplasia in preterm infant under ۳۰ week with Respiratory Distress Syndrome A Randomized Cilinical Trial Purpose Prevention Inclusion/Exclusion criteria Inclusion criteria: All preterm babies with birth age less than 30 weeks provided they have respiratory distress syndrome (RDS) There is evidence of RDS in chest X-ray and ABG and they are candidates to receive surfactant Babies born at the hip less than 24 weeks of pregnancy - with a birth weight of less than 400 grams - the presence of congenital cardiopulmonary disease - conditions for not entering the study Exclusion criteria: A baby born less than 24 weeks old A baby with a birth weight of less than 400 grams A baby with a fatal congenital abnormality or disorder Presence of cardiopulmonary diseases Age From 6 months old to 7 months old Gender Both Phase 3 Groups that have been masked No information Sample size Target sample size: 150 Randomization (investigator's opinion) Randomized Randomization description Randomization method: using block randomization with unequal block sizes (4 Two groups of comparison and intervention are interpreted. Confounding Control Method: Randomization of Allocation. Control in the analysis phase is different from statistical models. : Random allocation and concealment and blinding Hidden Privatization: By specifying unequal block sizes, the allocation sequence will be hidden. Pre-randomization list by web-provided software by the co-researcher who is in the interpretation process (https://www.sealedenvelope.com/simple-randomiser/v1/ists) He is not involved in preparing and only this researcher will have access to this list. At the time of identification of the qualified patient, the allocation group will be announced by him based on the prepared list. Blinding: The researcher evaluating the outcome of the specialized group of samples is known According to the fact that the patient is a baby and the parents will also perform the treatment without knowledge. The review of the final results of the study will be done by a doctor who was unaware of the type of intervention. For statistical analysis, a statistical expert is used who is not informed about how people are assigned to groups. Concealment Method Privatization Method: The problem allocation list is only available to the secretary of the department. In order to conceal the randomization process, the sequence of treatment is written on the	cards in order, then the cards are placed in sealed envelopes. Each envelope has a random 10-digit code without order and identification of the corresponding patient number, and the only design of the methodologist will be the corresponding code. Because the doctor declares the eligibility of a patient, the secretary provides the envelope to the doctor and the desired treatment is carried out according to the specific type mentioned in the envelope. Preterm infants who meet the conditions for entering the study in the form of available sampling and are placed in one of the following two groups based on the randomization list: mg/kg intervention in the first group including: intratracheal injection of surfactant 200 and followed by . mg/kg and surfactant 200 budesonide 0.25 mg/kg in the second intervention group including intratracheal injection. kg if needed Sampling Type: Available. Randomization method: Using block randomization with unequal block sizes (4-6-8), the samples are assigned to one of the comparison and intervention groups. Confounding Control Method: Randomization of Allocation. Control in the analysis phase is different from statistical models. Blinding (investigator's opinion) Not blinded Blinding description Placebo Not used Assignment Parallel Other design features Secondary Ids empty Ethics committees 1 Ethics committee Name of ethics committee ethics committee of Alborz University of Medical Sciences Street address Azimiyeh City Karaj Province Alborz Postal code 1463614636 Approval date 2022-09-24, 1401/07/02 Ethics committee reference number IR.ABZUMS.REC.1401.181 Health conditions studied 1 Description of health condition studied Bronchopulmonary dysplasia
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ICD-10 code

P27.1

ICD-10 code description

Bronchopulmonary dysplasia originating in the perinatal period

Primary outcomes**1****Description**

Severity of BPD: BPD will be divided into three groups in terms of severity: mild, which means that they do not need oxygen at the age of 36 weeks of pregnancy, and moderate, which means that they need less than 30% of oxygen in the 36th week of pregnancy. Severe need for 30% oxygen with or without positive pressure ventilation or continuous positive pressure at 36 weeks of gestation

Timepoint

After 28 days

Method of measurement

Physical examination

Secondary outcomes

empty

Intervention groups**1****Description**

Preterm infants who meet the conditions for entering the study are included in the study as available samples and are placed in one of the following two groups based on listrandomization: mg/kg intervention in the first group includes: intratracheal injection of surfactant 200 mg/kg

Category

Prevention

2**Description**

Control group: Control group: under sterile conditions, surfactant is prepared in a syringe for the control group, and for the intervention group: surfactant and budesonide are mixed in a syringe and gently shaken to obtain a uniform solution before injection. Medicines are injected intratracheally. In case of surfactant or with budesonide, it can be repeated every 8 hours until a maximum of 6 doses have been received. $FiO_2 < 0.3$, extubation when patients are hospitalized and not discharged, patients are visited daily, in the morning by the neonatologist, if they are discharged, they will be visited again after 72 hours.

Category

Prevention

Recruitment centers**1****Recruitment center****Name of recruitment center**

Kamali Hospital

Full name of responsible person

Sima Kianpishe

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Azimiye

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Sponsors / Funding sources**1****Sponsor****Name of organization / entity**

Karaj University of Medical Sciences

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Grant name**Grant code / Reference number****Is the source of funding the same sponsor organization/entity?**

Yes

Title of funding source

Karaj University of Medical Sciences

Proportion provided by this source

100

Public or private sector

Public

Domestic or foreign origin

Domestic

Category of foreign source of funding

empty

Country of origin**Type of organization providing the funding**

Academic

Person responsible for general inquiries

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Contact

Name of organization / entity
Karaj University of Medical Sciences
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Person responsible for updating data

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Sharing plan

Deidentified Individual Participant Data Set (IPD)

Undecided - It is not yet known if there will be a plan to make this available

Study Protocol

Undecided - It is not yet known if there will be a plan to make this available

Statistical Analysis Plan

Undecided - It is not yet known if there will be a plan to make this available

Informed Consent Form

Undecided - It is not yet known if there will be a plan to make this available

Clinical Study Report

Undecided - It is not yet known if there will be a plan to make this available

Analytic Code

Undecided - It is not yet known if there will be a plan to make this available

Data Dictionary

Undecided - It is not yet known if there will be a plan to make this available